ROZENFEL'D, I.L.; PAVIUTSKAYA, T.I.; FRUMKIN, A.N., akademik. New principle in the investigation of atmospheric corrosion of metals. (MLRA 6:6) Dokl. AN SSSR 91 no.2:315-317 J1 '53. 1. Akademiya nauk SSSR (for Frunkin). (Corrosion and anticorrosives)

FRUMKIN, A.N., akademik; MIYESSERCV, K.G.

Poisoning of aluminosilicate catalysts by pyridine bases. Dokl.AN SSSR 91 no.3:553-556 Jl '53. (MLRA 6:7)

1. Institut nefti Akademii nauk SSSR (for Miyesserov). 2. Akademiya nauk SSSR (for Frumkin). (Catalysts) (Alumimum silicates)

AXHUMOV, Ye.I.; SPIRO, N.S.; FRUMKIN, A.W., akademik.

Relation between concentrations of isoactive aqueous two-component solutions. Dokl.AN SSSR 91 no.3:573-576 J1 '53. (MLRA 6:7)

1. Akademiya nauk SSSR (for Frumkin). (Solutions (Chemistry))

FRUMKIN, A.H., akademik; FORSELOM, G.V.; MASHOVETS, V.P.

Conditions for the modeling of the electric field of electrolysers. Dokl.
AM SSSR 91 no.3:593-595 JI '55. (MLRA 6:7)

1. Vsesoyusnyy alyuminiyevc-magniyevyy institut (for Forsblom and Mashovets).
2. Akademiya nauk SSSR (for Frunkin).

(Electrolysis--Electromechanical analogies)

RYSS, I.G.; ML'KENBARD, A.G.; FRUMKIN, A.N., akademik.

Thermochemistry of fluo-boric complexes. Dokl.AN SSSR 91 no.4:865-868 Ag *53. (MIRA 6:8)

1. Akademiya namk SSSE (for Frunkin). 2. Depropetrovskiy metallurgi-cheskiy institut im. I.V.Stalina (for Ryss and Elkenbard). 3. Depropetrovskiy sel'skokhozyaystvennyy institut (for Ryss and Elkenbard). (Compounds, Complex) (Boron) (Fluorine)

BELYAYEV, I.N.; BERGMAN, A.G.; NOMIKOS, L.I.; FRUMKIN, A.N., akademik.

Electric conductivity of diagonal cross sections of ternary reciprocal system in melts. Dokl.AN SSSR 91 no.5:1103-1105 Ag '53. (MLRA 6:8)

1. Akademiya nauk SSSR (for Frumkin). 2. Rostovskiy-na-Donu gosudarstvennyy universitet (for Belyayev, Bergman, Nomikos).

(Phase rule and equilibrium) (Electric conductivity)

BLOKH, G.A.; FRUMKIN, A.N., akademik.

Investigation of the mechanism of action of accelerators for rubber vulcanization. Vulcanization of rubber with sulfur isotopes. Dokl.AN SSSR 91 no. 5:1107-1110 Ag '53. (MLRA 6:8)

1. Akademiya nauk SSSR (for Frumkin). 2. Kievskiy tekhnologicheskiy institut legkoy promyshlennosti. (Vulcanization) (Sulfur--Isotopes)

ISAGULYANTS, G.V.; ANDREYEV, Ye.A.; KOSOLAPOVA, N.A.; FRUMKIN, A.N., akademik.

Obtaining caproic acid tagged with radiocarbon, C¹⁴, in carboxyl. Dokl.AN SSSR 91 no.5:1123-1124 Ag '53. (MLRA 6:8)

1. Akademiya nauk SSSR (for Frumkin). (Caproic acid) (Carbon—Isotopes)

APPROVED FOR RELEASE: 06/13/2000 CIA-RDP86-00513R000513820007-5"

HAT THE TANK

IOFA, Z.A.; ROZHDESTVENSKAYA, G.B.; FRUMKIN, A.N., akademik.

Measuring the adsorption of iedine ions on iron. Dokl.AN SSSE 91 no.5;1159-1162 Ag 153. (MLRA 6:8)

1. Akademiya nauk SSSR (for Frunkin).
(Adsorption) (Iedine) (Iron)

ZADUMKIN, S.N.; FRUMKIN, A.N., akademik.

Surface tension and heat of evaporation of metals. Dokl.AN SSSR 92 no.1: 115-118 S '53. (MLRA 6:8)

1. Akademiya nauk SSSR (for Frumkin). 2. Kabardinskiy gosudarstvennyy pedagogicheskiy institut (for Zadumkin). (Metals)

GOLIDER, G.A.; ZHDANOV, G.S.; UNANSKIY, M.H.; FRUMKIN, A.N., akademik.

Radiographic determination of the structure of picryl chloride. Dokl.AN SSSR 92 no.2:311-314 S '53.

(HLRA 6:9)

1. Akademiya nauk SSSR (for Frumkin). 2. Fiziko-khimicheskiy institut im. L.Ya.Karpova (for Gol'der. Zhdanov and Umanskiy). (Radiography) (Picryl chloride)

BORESKOV, G.K.; SLIN'KO, M.G.; FILIPPOVA, A.G.; FRUMKIN, A.H., akademik.

Catalytic activness of nickel, palladium, platinum, in respect to the reaction of oxidating hydrogen. Dokl.AN SSSR 92 no.2:353 S '53. (MLRA 6:9)

1. Akademiya nauk SSSR (for Frumkin).

(Oxidation) (Hydrogen) (Catalysts)

FIALKOV, Ya.A.; BUR'YANOV, Ya.B.; FRUMKIN, A.N., akademik.

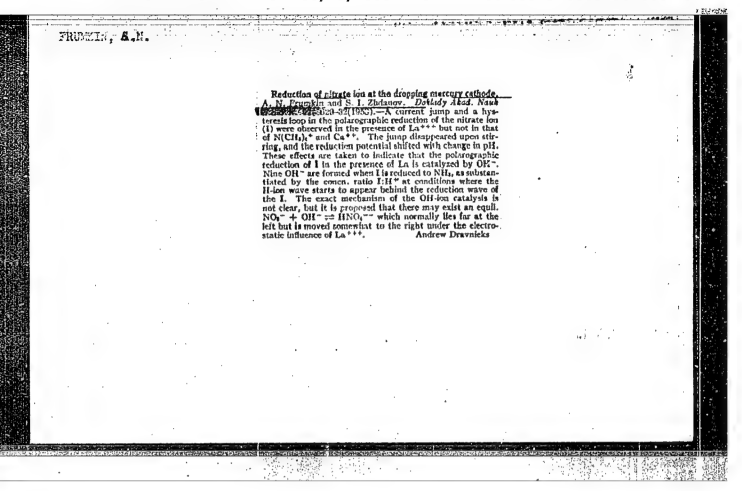
Complex compounds of PCl₅ with aluminum and iron chlorides. Dokl.AN SSSR 92 no.3:585-588 S 153. (MLRA 6:9)

1. Akademiya nauk SSSR (for Frumkin). 2. Institut obhschey i neorganicheskoy khimii Akademii nauk Ukrainskoy SSR .
(Compounds, Complex) (Chlorides)

ZAYDES, A.L.; STOYAHOVA-SINITSKAYA, 1.G.; FRUMKIN, A.H., akademik.

Ordered structure of films of hydrated cellulose. Dokl.AN SSSR 92 no.3: 601-602b S 153. (MLRA 6:9)

1. Akademiya nauk SSSR (for Frumkin). 2. TSentral'nyy nauchno-issledova-tel'skiy institut kozhevenno-obuvnoy promyshlennosti (for Zaydes and Stoyanova-Sinitskaya). (Cellulose)



ZHDANOV, G.S.; ZHURAVLEV, N.N.; ZEVIN, L.S.; FRUMKIN, A.N., akademik.

Radiographic determination of the formation of solid solutions in baron carbide. Dokl.AN SSSR 92 no.4:767-768 0 153. (MIRA 6:9)

1. Akademiya nauk SSSR (for Trumkin). 2. Moskovskiy mekhanicheskiy institut Ministerstva kul'tury SSSR (for Zhdanov, Zhuravlev and Zevin)
(Baron carbide) (Radiography)

RYABCHIKOV, D.I.; LAZAREV, A.I.; FRUMKIN, A.N., akademik.

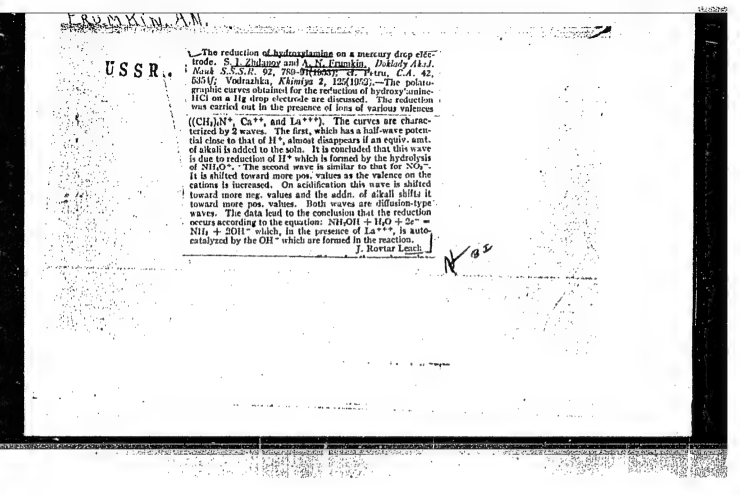
Chromatographic separation of molybdenum and rhenium. Dokl.AN SSSH 92 no.4: 777-779 0 '53. (MLRA 6:9)

1. Akademiya nauk SSSR (for Frumkin). 2. Institut geokhimii i analiticheskoy khimii im. V.I. Vernadskogo Akademii nauk SSSR (for Ryabchikov and Lazarev).

(Chromatographic analysis) (Molybdemum) (Rhenium)

"APPROVED FOR RELEASE: 06/13/2000

CIA-RDP86-00513R000513820007-5



UKSHE, Ye.A.; LEVIN, A.I.; FRUMKIN, A.N., akademik.

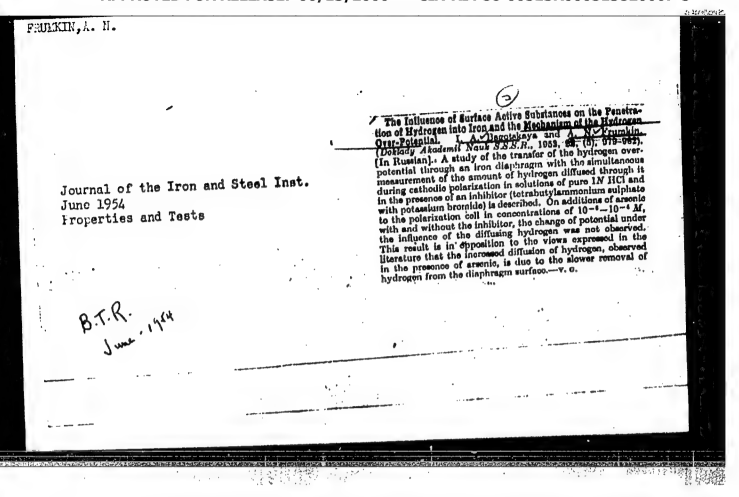
Effect of the adsorption of cations on electrodeposition of copper from pyrophosphate electrolytes. Dokl. AN SSSR 92 no.4:799-801 0 153. (MLHA 6:9)

1. Akademiya nauk SSSR (for Frumkin). 2. Ural'skiy politekhnicheskiy institut im. S.M.Kirova, Sverdlovsk (for Ukshe and Levin).

(Copper--Electrometallurgy) (Pyrophosphates)

"APPROVED FOR RELEASE: 06/13/2000

CIA-RDP86-00513R000513820007-5



KUZNETSOV, V.A.; KOCHERGIN, V.P.; TISHCHENKO, M.V.; POZDNYSHEVA, Ye.G.; FRUMKIN, A.H., akadenik.

Investigation of surfaces tension of the alloy: tin - cadmium on the boundary with the fused eutectic: Li - KCl in a vacuum. Dokl.AN SSSR 92 no.6:1197-1199 0 '53.

1. Akademiya nauk SSSR (for Frumkin). 2. Ural'skiy gosudarstvennyy universitet im. A.M.Gor'kogo, gorod Sverdlovsk (for Kusnetsov, Kochergin, Tishchenko (Cadmium-tin alloys) (Surface tension) and Posdnysheva).

ZHDANOV, G.S.; POSPELOV, V.A.; PRUMKIN, A.N., akademik.

Unequal length of metal oxygen bonds in certain metal oxides and the "molecular" structure ZnO. Dokl.AN SSSR 93 no.1:97-99 N '53. (HURA 6:10)

1. Akademiya nauk SSSR (for Frumkin). 2. Fiziko-khimicheskiy institut im. L. Ya. Karpova (for Zhdanov and Pospelov). (Metallic oxides)

APPROVED FOR RELEASE: 06/13/2000 CIA-RDP86-00513R000513820007-5"

一种主要的数据不在独立

NOVOKHATSKIY, I.P.; KALININ, S.K.; FRUNKIN, A.N., akademik.

Fluorine content of certain natural waters of Kasakhstan according to spectral analysis data. Dokl.AN SSSR 93 no.2:289-291 N '53. (MLRA 6:10)

1. Akademiya nauk Kaz.SSR, Alma-Ata. 2. Akademiya nauk SSSR (for Frumkin).

(Kazakhatan--Water--Analysis) (Analysis--Water--Kazakhatan)

(Fluorine)

KACHURIN, L.G.; FRUMKIN, A.N., akademik.

Probability of the formation of ice nuclei in supercooled water. Dokl.AN SSSR 93 no.2:307-310 N 153. (MLRA 6:10)

1. Leningradskiy gidrometeorologicheskiy institut (for Kachurin). 2. Akademiya nauk SSSR (for Frumkin). (Ice)

D'YAKOV, A.A.; FRUMKIN, A.N., akademik.

Certain problems in the electroreduction of persulfate. Dokl.AH SSSR 93 no. 4:685-687 D '53. (MLRA 6:11)

1. Akademiya nauk SSSR (for Frunkin).
(Reduction, Electrolytic) (Persulfates)

SAMSONOV, G.V.; FRUMKIN, A.N., akademik.

Physical properties of certain intrusion phases. Dokl.AN SSSR 93 no.4:689-

692 D '53. (MERA 6:11)

1. Akademiya nauk SSSR (for Frumkin). 2. Moskovskiy institut tsvetnykh metallov i solota im. M.I.Kalinina (for Samsonov).

(Metals) (Solutions, Solid)

FRUMKIN, A.N.

USSR/Chemistry - Physical chemistry

Card 1/1

Pub. 147 - 14/26

Authors

Mayzlish, R. S.; Tverdovskiy, I. P.; and Frumkin, A. N.

Title

Adsorption phenomena on Hg and alcohol-water solutions of electrolytes

Periodical

Zhur. fiz. khim. 28/1, 87-101, Jan 1954

Abstract

Two series of electro-capillary curves representing ethyl alcohol - water - NaCl(sat) and ethyl alcohol - water NH₄Cl(sat) systems were measured and it became evident that the boundary tension of the aqueous salt solution decreases in the zone of electro-capillary maximum as result of introduction of alcohol into the solution. Addition of water to the alcohol salt solution leads to reduction of boundary tension on both ends of the electro-capillary curve. It was established that the negative adsorption of alcohol corresponds to the positive adsorption of water. The calculated adsorption isotherms on the boundary with air showed that both salts increased the alcohol adsorption and displace the adsorption maximum toward less concentrated alcohol solutions. Ten references: 5-USSR; 4-USA and 1-English (1913-1953). Tables; graphs.

Institution :

Institute of Applied Chemistry, Leningrad

Submitted:

March 14, 1953

June 11, 1953

Submitted

FRUMKIN, A. N USSR/Chemistry	•		
Card 1/1			
Authors		Kalish, T. V., and Frumkin, A. N.	
Title	1	Effect of anion adsorption on the reduction of an S208-ion on a drop cathode	
Periodical		Zhur. Fiz. Khim. 28, Ed. 3, 473-489, March 1954	
Abstract	:	It is shown that the reaction of electro-reduction of an S208-Ion on a mercury electrode is retarded by the adsorbing anions. The rate of this reaction, at a positive surface charge in relation to the nature of the indifferent electrolyte, decreases gradually. The potential at which the current intensity corresponding to this reaction reaches 0.8 in proportion to the maximum diffusion current is displaced toward the negative side. At a negative surface charge one can observe a certain weakly expressed dependence of the rate of electro-reduction upon the nature of the anions which in this case are oriented in reverse sequence. The effect of adsorption anions leads to the convergence of the potential with the point of the zero-charge. Thirty references; 1 USSR. since 1919. Tables, graphs.	The state of the s
Institution		Acad. of Sc. USSM Institute of Physical Chemistry, Moscow	

FRUNKIN, AN. TRUSKIN, A. N. USSR/Chemistry

Card 1/1

Authors

: Kalish, T. V., and Frunkin, A. N.

Title

: Position of the zero-charge point and its effect on the electro-reduction of a S20g anion on a drop electrode

Periodical

: Zhur. Fiz. Khim., 28, Ed. 5, 801 - 805, May 1954

Abstract

: Experiments showed that, the beginning of the descent of the polarization curve, during electro-reduction of S20g on a drop electrode consisting of 40% thallium amalgam, is displaced by 0.5 v toward more negative potentials, which corresponds with the position of the zero charge point of thallium amalgam. The retardation of the S208 reduction reaction over a mercury electrode in solutions containing n-octyl alcohol is determined by the absorption of alcohol molecules, the action of which is affected by the negative charges of the surface. Seven USSR references. Graphs.

Institution : Acad. of Sc. USSR, Institute of Physical Chemistry, Moscow.

Submitted

: July 8, 1953

FRUMKIN, A. N.

USSR/Physics

Card 1/1

Frumkin, A. N., Academician, and Zhdanov, S. I. Authors

Origination of sudden changes in current intensity and Title

hysteresis loop on lolarization curves during electro-re-

duction reaction

Dokl. AN SSSR, 96, Ed. 4, 793 - 796, June 1954 Periodical

During the reduction of NO3 and NO2 ions on a mercury drop electrode in the presence of La3+, an intermittent increase Abstract

in the current intensity may be observed when a certain potential is reached. The anomalous form of the polarization curves is due to the autocatalytic effect of OH, formed during the reduction of ions. An approximate theory of these phenomena is given. The hysteresis loop is best expressed at PH = 3 - 4 and disappears at a much higher acidification and alkalization.

and alkalization. Three references. Graphs.

Institution :

April 7, 1954 Submitted

"APPROVED FOR RELEASE: 06/13/2000

CIA-RDP86-00513R000513820007-5

FRUMIN, A. N.

USSR/Chemistry - Physical chemistry

Card 1/1

: Pub. 22 - 29/48

Authors

. : Frumkin, A. N., Academician, and Zhdanov, S. I.

Title

* Reduction of a nitrate-ion in an acid medium in the presence of La

Periodical

1 Dok. AN SSSR 97/5, 867-870, August 11, 1954

Abstract

The phenomena observed during the reduction of a nitrate-ion, i. e., in conditions when the reaction of the solution in the near-electrode layer cannot transform into the alkali zone because of the continuing reaction process, are discussed. The reduction of a NO3 ion in an acid medium is followed by the reaction of H-separation. The final product of NO₃ reduction (ammonia), at pH values attainable in the presence of La3+ surplus, is included in the solution, basically in the form of a NH₄ - ion. Three references: 2-USSR and 1-Polish

(1951-1954). Graphs.

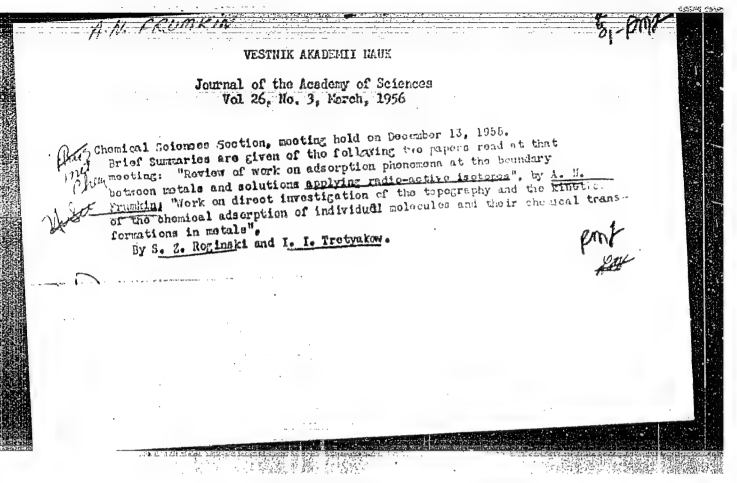
Institution

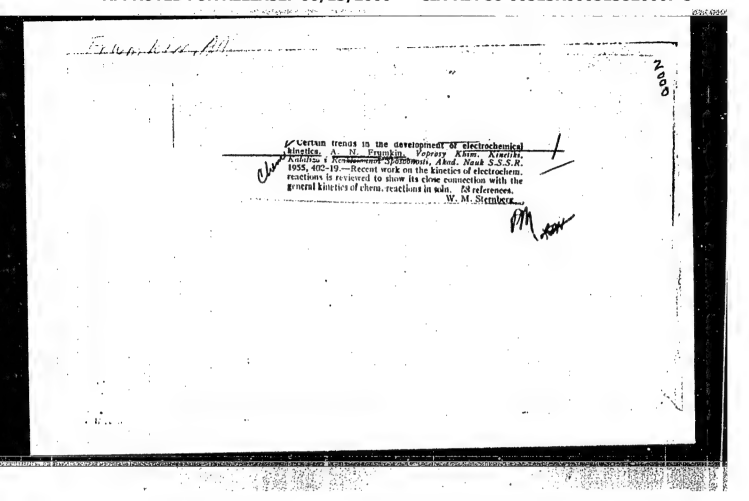
Submitted

June 16, 1954

"APPROVED FOR RELEASE: 06/13/2000

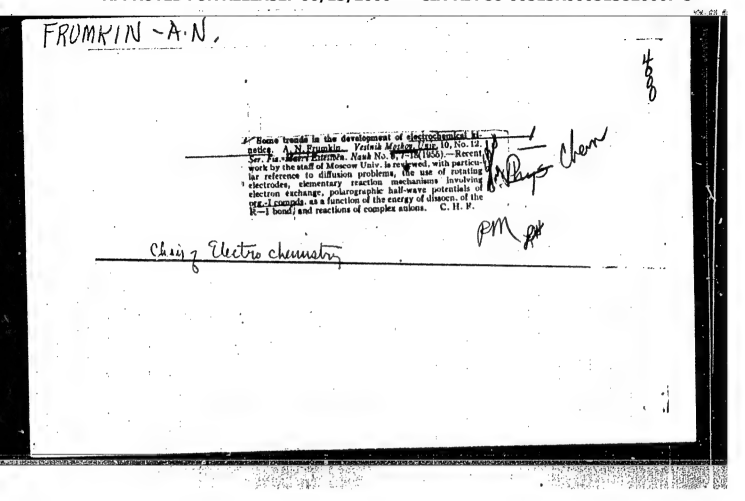
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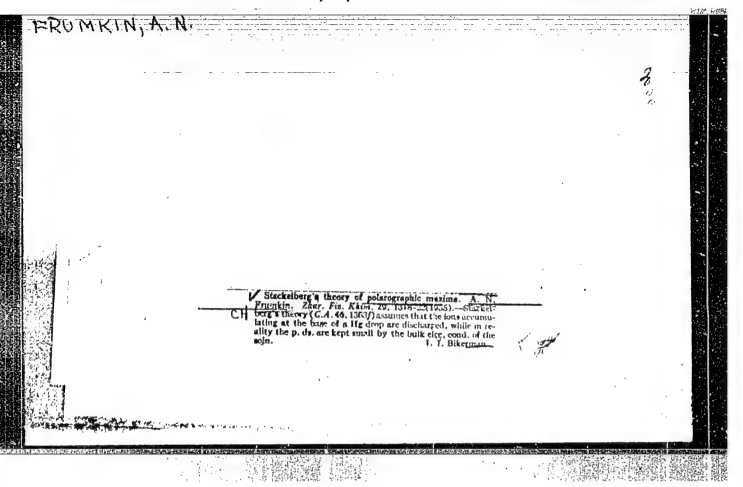


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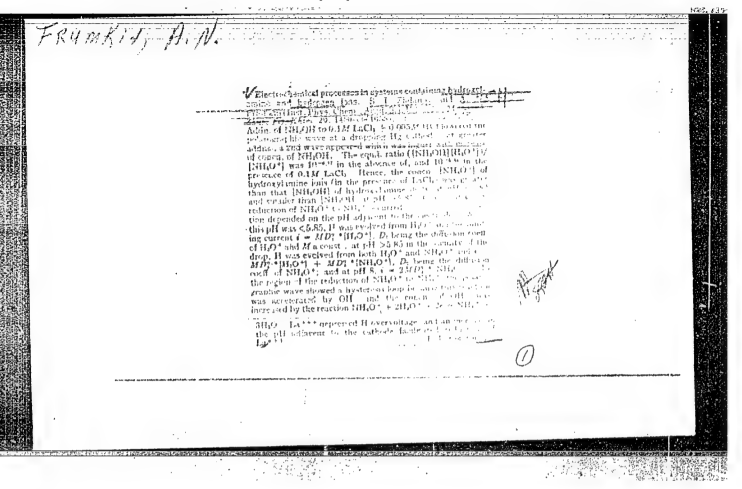


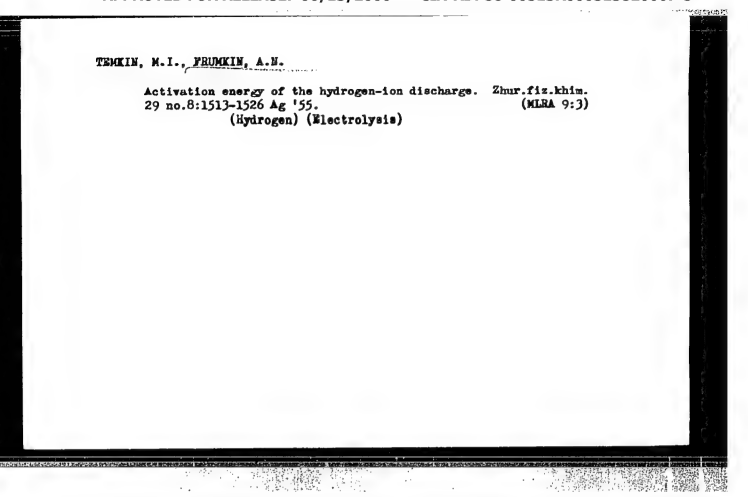
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	(Russian.) A. N. Frum p. 933-950.	skin. Uspekhi khimil, v. 24, no. 8, 193	s. Netal	
	Relation of alternating external friction of meta-	a current canacitance, hardness, as al to the potential of a metallic electro- tables. 82 ref.	de /	
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"APPROVED FOR RELEASE: 06/13/2000

CIA-RDP86-00513R000513820007-5





FRUMKIN, A.N.

*USER/ Chemistry - Physical chemistry

Card 1/2

Pub. 147 - 11/21

Authors

: Florianovich, G. M., and Frumkin, A. N.

Title

Electro-reduction of anions over a mercury electrode

Periodical

Zhur. fiz. khim. 29/10, 1827-1846, Oct 1955

Abstract

The reduction of numerous anions was investigated on a mercury drop and amalgamated rotating cathodes. It was found that anions begin reducing at sufficiently negative values of the potential when the electrode surface is charged negatively (relatively difficult reducible anions) and that the anion reduction begins at such potentials at which the electrode surface is positively charged (easily reducible anions). The effect of foreign cations

Institution:

Moscow State University im. M. V. Lomonosov

Submitted

: February 15, 1955

"APPROVED FOR RELEASE: 06/13/2000

CIA-RDP86-00513R000513820007-5

Card 2/2

Pub. 147 - 11/21

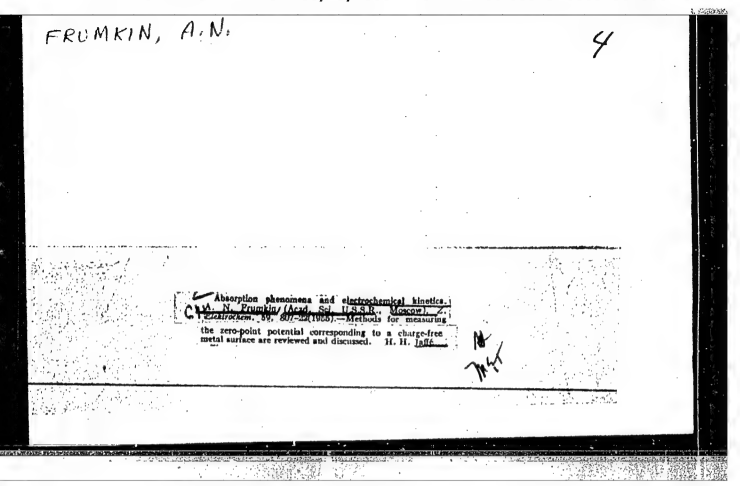
Periodical

Zhur. fiz. khim. 29/10, 1827-1846, Oct 1955

Abstract

on the anion reducing process was investigated and it was established that the cations affect the magnitudes of minimum currents and increase same. The effectiveness of the cations depends upon their charge and concentration. Thirty-four references: 7 Czech., 6 USA, 18 USSR, 1 Fr.

and 2 Eng. (1932-1954). Table; graphs; drawing.



FRUMKIN, A.N.

USSR/ Chemistry - Physical chemistry

Card 1/2

Pub. 22 - 31/52

Authors

Frunkin, A. N. Academician., and Aykazyan, E. A.

Title

The kinetics of ionization of molecular hydrogen on a Pt electrode and

the role of anions

Periodice

Dok. AN SSSR 100/2, 315-318, Jan 11, 1955

Abstract

Investigation was conducted to determine the relation between the current density of molecular hydrogen ionization on a disc-type smooth Pt electrode and the potential and nature of the anion at various rpas. The hydrogen dissolved was kept in equilibrium with the hydrogen at

atmospheric pressure.

Institution :

The M. V. Lomonosov State University, Moscow

Submitted

October 8, 1954

Periodical :

Dok. AN SSSR 100/2, 315-318, Jan 11, 1955

Card 2/2

Pub. 22 - 31/52

Abstract

It was found that the processes of ionization of adsorbed H and discharge of H ions are practically in reverse order and that the rate of the process on the whole is determined by the hydrogen adsorption and desorption stages. The effect of the anions was found to be the strongest when the electrode potentials were most positive. Ten references: 9 USSR and 1 German (1935-1952). Graphs.

Frum Kin, A.N.

USSR/Chemistry - Electrochemistry

Card 1/1

Pub. 22 - 34/54

Authors

Frunkin, A. N., Academician; Kaganovich, R. I.; Gerovich, M. A.; and

Vasilyev, v. R.

Title

The mechanism of anodic formation of persulfates

Periodical : Dok. AN SSSR 102/5, 981-983, Jun 11, 1955

Abstract

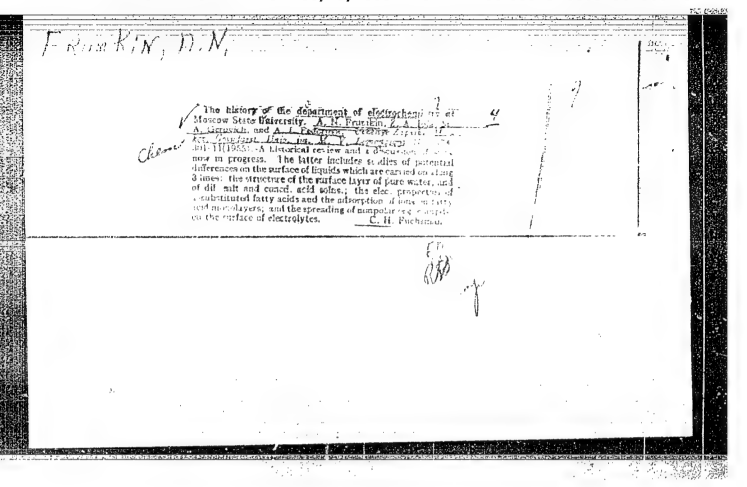
Sulfate electrolysis experiments were carried out in water enriched with the heavy 018 isotope, in alkali and weak acid electrolyte at possibly low temperatures to establish the condition most favorable for the anodic formation of persulfates. The results indicate that the first product formed on the anode is an oxygen-containing water-oxidation compound OH which in turn oxidizes the SOZ- ion within the volume of the solution. Seven references: 3 USSR, 3 USA and 1 German (1922-1954). Table.

Institution : Acad. of Sc., USSR, Inst. of Phys. Chem. and the M.V. Lomonosov State

University, Moscow

Submitted

April 11, 1955



USSR/General Problems - Methodology. Scientific Institutions A-1 and Conferences. Instruction. Questions Concerning Bibliography and Scientific Documentation.

Abs Jour : A.N. Frumkin.

A 6

Inst : Title : In Memory of a Scientist and Revolutionist.

Orig Pub : Khim. nauka i prom-st', 1956, 1, No 5, 573-574

Abstract : To the 10th anniversary of the death of A.N. Bakh

(1857 - 1946).

Card 1/1

TRE 4:11

_ 4 _

TEMKIN, M.I.; PRUMKIN, A.N.

On the paper by O.M.Pelterak "The activation energy of hydrogen ion discharge and the problem of absolute potential in electrochemical kinetics". Zhur.fis.khim.30 no.5:1162-1168 My '56. (MIRA 9:9)

(Hydrogen) (Petential, Theory of) (Pelterak, O.M.)

Potential difference at the water - air interface [with English summery in insert]. Zhur.fiz.khim. 30 no 7:1455-1468 J1 '56.

(MLRA 9:11)

1. Moskovskiy gosudarstvennyy universitet imeni M.V.Lomonosova.

(Electrochemistry) (Potential, Theory of)

FRUMKIN, A.N.

Category: USSR / Physical Chemistry - Electrochemistry.

B-12

Abs Jour: Referat Zhur-Khimiya, No 9, 1957, 30132

Author : Temkin M. I., Frumkin A. N.

: Comments on the Dependence of Hydrogen Overvoltage on Nature of Title

the Cathode in Connection with the Paper by Ryuchi and Delae

(Ryutshi and Delakhey in the Original).

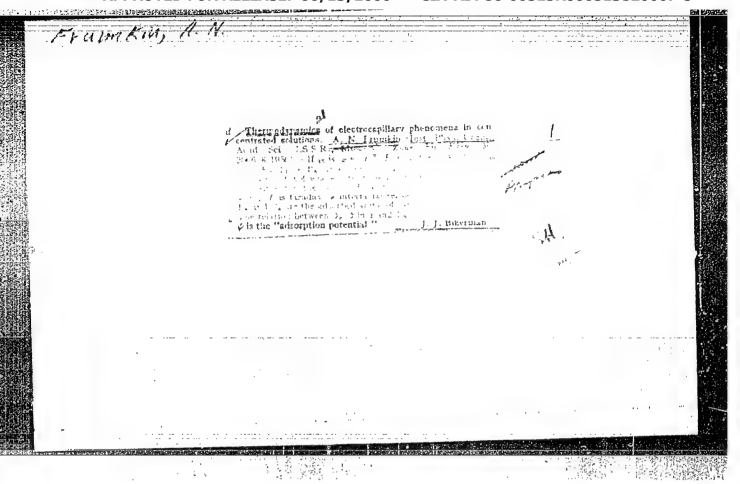
Crig Pub: Zh. fiz. khimii, 1956, 30, No 8, 1885-1988

Abstract: A discussion article (RZhKhim, 1956, 12527); see also RZhKhim, 1956,

54066.

: 1/1 Card

-10-



"APPROVED FOR RELEASE: 06/13/2000

CIA-RDP86-00513R000513820007-5

"A Contribution to the Overvoltage Theory of Hydrogen," Zeltschrift Fuer Physikalische Chemie," (Periodical of Physical Chemistry), October 1957.

YAKOBI, B.S.; YRUMKIN, A.M., akademik, red.; SMIRNOVA, A.V., tekhn.red.

[Work on electochemistry; a collection of articles and oapers]
Raboty po elektorkhimi; abornik state1 i materialov. Pod red.
Raboty po elektorkhimi; abornik state1 in materialov. Pod red.
A.M.Frumkina. Moskva, Ind-vo Akad.uauk SSSR, 1957. 301 p.

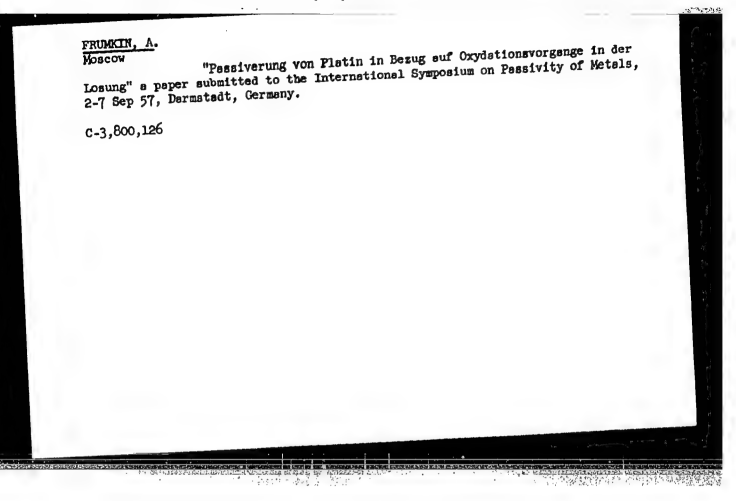
(Electrochemistry)

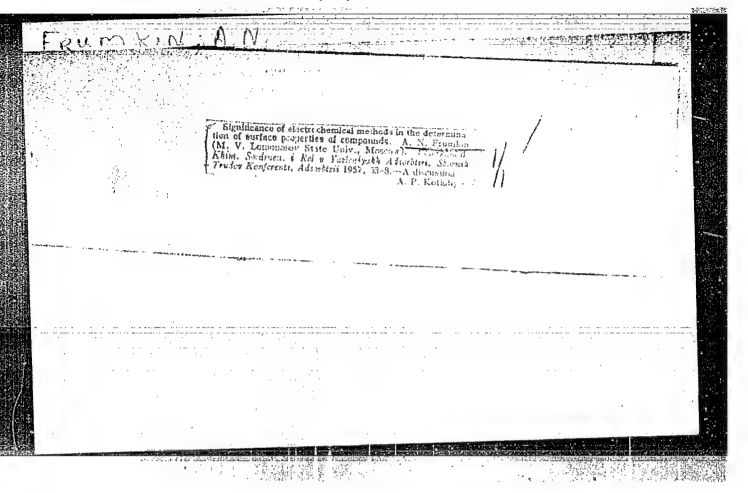
(MIRA 11:2)

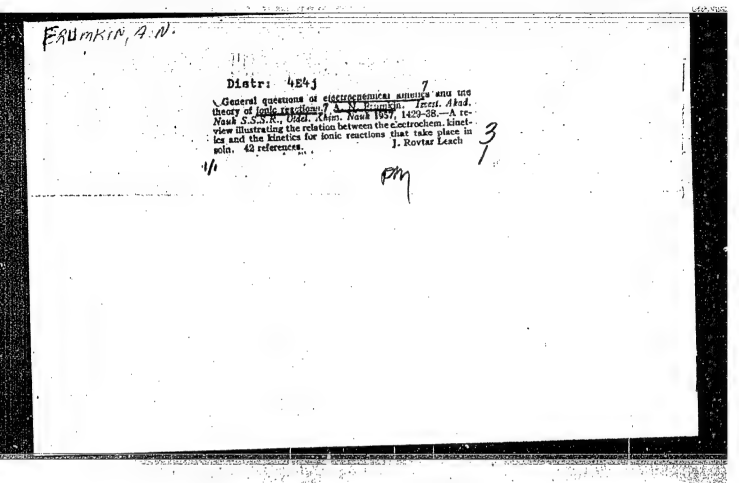
FRUMKIN, A. N.

Adsorption of organi substances at the metal electrolyte solution interface and its influence on electrochemical processes. Academy of Science, USSR, Physical Chemical Laboratory of Surface Phenomena, Moscow, U.S.S.R.

A paper to be presented at the 2nd International Congress of Jurface Activity London, April 1957

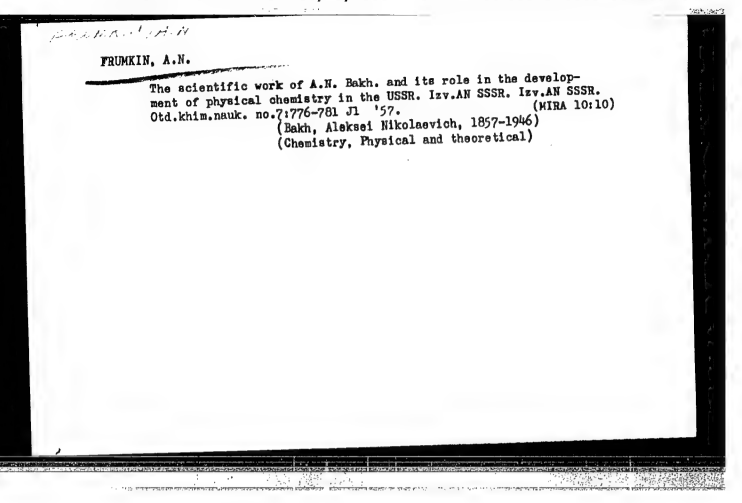






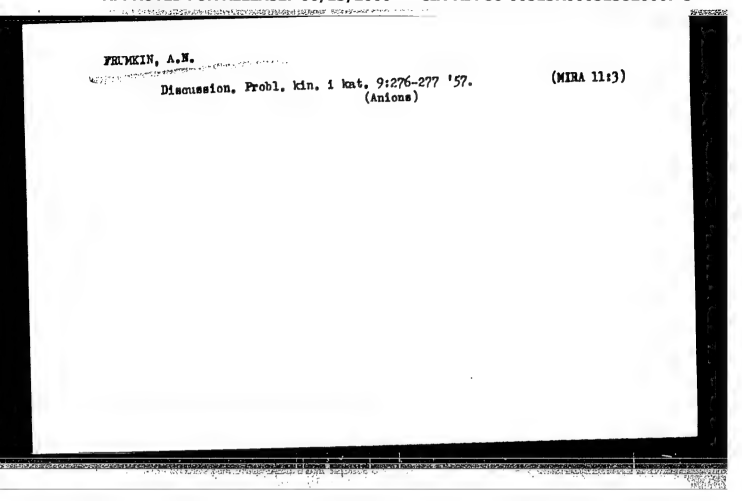
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Bunsay AN

AUTHOR:

Frumkin, A. N., Academician

30-10-13/26

TITLE:

Tasks of the Institute of Electrochemistry

(Zadachi instituta elektrokhimii).

PERIODICAL: Vestnik AN SSSR, 1957, October, Nr 10, pp. 99-101 (USSR)

ABSTRACT:

The electrochemistry has to solve a series of great tasks within the sixth Five-year Plan. therefore to organize within its scope proper institutes of

The problems set to these institutes can be characterized

1) Enlarging of the theoretical electrochemistry in order to clarify especially the question of electrochemical kinetics and thermodynamics.

2) Obtaining of new chemical sources of current which satisfy severest service conditions without relaxing.

3) Obtaining of new secon:dary electrochemical sources of current from atomic energy.

4) Determination of the phenomena occuring between semiconductors and electrolytic solutions.

5) Clarification of the processes at the primary and secundary electron emission correlated with the presence

Card 1/2

30-10-13/26

Tasks of the Institute of Electrochemistry

of contact potentials at the limits between metals and semi-conductors.

6) The possibilities of electrolysis on a large technical scale should be studied in connection with the construction of new large electric power plants.

Library of Congress AVAILABLE:

Card 2/2

FRUMKIN, A.H.; HIKOLAYEVA-FEDOROVICH, N.V.

Electric reduction of anions and adsorption of cations. Vest. Mosk.
un. Ser. mat.mekh. astron. fiz. khim. 12 no.4:169-184 '57.
un. Ser. mat.mekh. astron. fiz. khim. 12 no.4:169-184 (MRA 11:5)

1.Kafedra elektrokhimii Moskovskogo gosudarstvennego universiteta.

(Electrochemistry) (Anions) (Cations)

"APPROVED FOR RELEASE: 06/13/2000

CIA-RDP86-00513R000513820007-5

62-12-2/20 Frumkin, A.N. AUTHOR: On Some General Problems of the Electrochemical Kinetics and the Theory of Ion Reactions (O nekotorykh obshchikh voprosakh elektro-TITLE: khimicheskoy kinetiki i teorii iomykh reaktsiy) Report Delivered at the Meeting of the Chemical Department of the AN USSR on October 31, 1957 (Doklad na sessii otdeleniya khimicheskikh nauk Akademii nauk SSSR, 31 oktyabrya 1957 g). Izvestiya AN SSSR Otdeleniye Khimicheskikh Nauk, 1957, Nr 12, PERIODICAL: pp. 1429-1438 (USSR) The reviewer distinguishes between three stages in the development of modern chemistry; in his report, however, he deals with the last ABSTRACT: stage, i.e. the development of electrochemical kinetics. The basic condition was to penetrate more deeply into the metal-electrolyte boundary structure. The results achieved are described as very modest. It is further said that a number of new physical methods of investigation are applied. The author then deals with the adsorption of gases on the surface of solid metals. The determination of the dependence of the velocity of the heterogeneous chemical process on the pressure of the gases taking part in the reaction requires knowledge of the isothermal lines of adsorption of the surfaces of the catalyst. Card 1/3

62-12-2/20

On Some General Problems of the Electrochemical Kinetics and the Theory of Ion Reactions. Report Delivered at the Meeting of the Chemical Department of the AN USSR on October 31, 1957

For this purpose in most cases the isothermal lines of adscrption according to Lengayur (Languair) were used. Measurements, which were carried out last by Shlygin, Ershler and the author between 1929 and 1938, led to the unexpected conclusion that the quantity of hydrogen adsorbed on platinum is linearly modified in first approximation (see formulas p. 1431). The author then deals with the electric capillary phenomena and anomalous properties of thin layers of liquids. After describing the hitherto distinguished 2 kinds of transition (from the adsorbed layer to the new phase), the author mentions the results obtained recently by Dyryagin and his collaborators: he was able, by means of optical measurements, to show that by approximation to the point of saturation the strength of the adsorbed layers of polar liquids (as e.g. of water and of alcohol) on the smooth surface of glass actually attains some dozen molecules (up to 100 A). The author dealt in detail with the result of the investigation of the ratio between the electrode processes and the ion reactions in solvents (see fig. 1). Independent of the conception formed of the participation of the adsorbed anions in the elementary

Card 2/3

On Some General Problems of the Electrochemical Kinetics and the Theory of Ion Reactions. Report Delivered at the Meeting of the Chemical Department of the AN USSR on October 31, 1957

62-12-2/20

process, there is no doubt that they form little bridges (mostiki), which connect the electrode with the cation [23]. There are 3 figures and 42 references, 33 of which are Slavic.

ASSOCIATION:

Institute for Physical Chemistry of the AN USSR (Institut

fizicheskoy khimii Akademii nauk SSSR).

SUBMITTED:

October 1, 1957

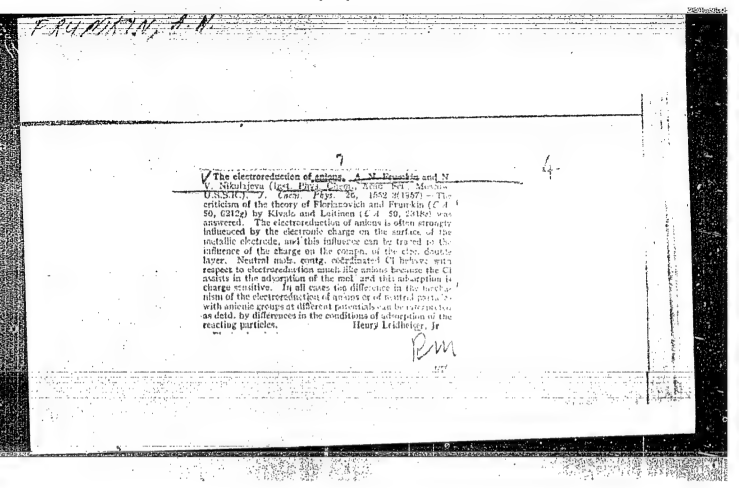
AVAILABLE:

Library of Congress

Card 3/3

L. Chemical engineering-Conference 2. Gas-Adsorption-Metals

3. Electrochemical kinetics



"APPROVED FOR RELEASE: 06/13/2000

CIA-RDP86-00513R000513820007-5

Tasks of the Institute of Electrochemistry. Vest. AN SSSR [27]

10.10:99-101 0 '57.
(Electrochemistry-Research)

"APPROVED FOR RELEASE: 06/13/2000 (

CIA-RDP86-00513R000513820007-5

FRUMKIN, A.P.

CONTRACTOR DE LA CONTRA

Frumkin, A. N., Member of the Academy.

30-11-18/23

AUTHOR:

Meetings with Polish and German Chemists (Vstrechi s pol'skimi i nemetskimi khimikami)

TITLE:

Vestnik AN SSSR, 1957, Vol. 27, Nr 11, pp. 122-128 (USSR)

ABSTRACT:

PERIODICAL:

After Polish electrochemists had paid a visit to the Institute for Physical Chemistry AN USSR the author stayed in Warsaw as a guest of the Polish Academy of Science. The research worksin the field of electrochemistry are concentrated in the Institute for Physical Chemistry under the direction of the eminent scientist V. Sventoslavskiy. The Smyalovskiy-Laboratory (physical chemistry of electrode-processes) is on a very high scienitific level; the investigations of the dependence of the penetration of hydrogen on the composition of various admixtures in steel, such as oxygen, sulphur, phosphor, are carried out on a broad basis. The collaborator B. Baranovskiy succeeded in discovering some new effects of motion of the liquid upon the passage of current. Pure calculation works on various problems of quantum-chemistry are done in the Laboratory S.Mintsa by theoretical physicists (the students of the well-known physicist L. Infel'd). The Laboratory for Physico-Chemical Methods of Analysis (under the direction of

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Meetings with Polish and German Chemists.

30-11-18/23

V.Kemul') which is connected with the chair for anorganic analytical chemistry of the Warsaw University, is distinguished by a special many-sidedness and topicality of themes. A method for determining minute traces of admixtures in metals was worked out in this laboratory. Much work is also deveted to the investigations of the mechanism of electrode-processes. Thus, for example, a marked breaking of the electrode-processes by absorbed foils of the hydroxidation was determined. In the Laboratory B. Kamenskogo the investigation of the potentials on the free surface of various solutions is successfully performed. The aim of these works is a comparison of these potentials with the physiological activity of organic compounds. Many works are distinguished by special originality. -Then the author reports on his impressions during the visit of the scientific research institutes in the DDR (GDR). He participated in the Berlin Colloquy on the kinetics of electrode-processes. K.Shvabe (Dresden), chairman of the organizationcommittee, talked on the theory of the overvoltage of hydrogen. M.Brayter (Munich) reported on the investigation of the anodeformation and the cathode-restoration of the surface-oxidations on electrodes of noble metals in acid solutions. Of special interest for the author was the detailed exchange of opinions in a

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Meetings with Polish and German Chemists.

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small circle on the most important still unsolved problems of electrochemical kinetics. In the technical college in Dresden the author held a lecture on "The restoration of anions in an electrical manner and the adsorption of cations" and in the Academia Leopoldina in Halle he talked on "The adsorption of organic substances on the boundary metal-solution and its influence upon the electrochemical processes."

AVAILABLE:

Library of Congress

Card 3/3

USSR/Physical Chemistry - Electrochemistry

B-12

Abs Jour

: Referat Zhur - Khimiya, No 1, 1958, 560

Author

A.N. Frumkin, A.S. Titiyevskaya.

Inst

Parameter and parameters .

Title

Electrocapillary Phenomena in Solutions of Thallium Salts.

Orig Pub

Zh. fiz. khimii, 1957, 31, No 2, 485-492

Abstract

: Electrocapillary curves (EC) for the solutions 1 n. H₂SO₄ + x n. Tl₂SO₄ (x = 0.01; 0.05; 0.1 n.) and 1 n. KHO₃ + + 0.01 n HNO₃ + x n. TlNO₃ (x = 0.01; 0.1 and 0.2 n.) were plotted in the region of the polarization of the Hg electrode from +0.1 to -0.45 v (satur. c. e.). A drop of the boundary tension was revealed at the maxima of the curves, as well as a shift of these maxima in the positive direction, which reached 0.29 v in the case of the solution 0.2 n. RlNO₃. A little surface activity of Tl⁺ ions was revealed also at positive charges of the Hg surface. The conclusion was made that Tl⁺ behaves as a typical

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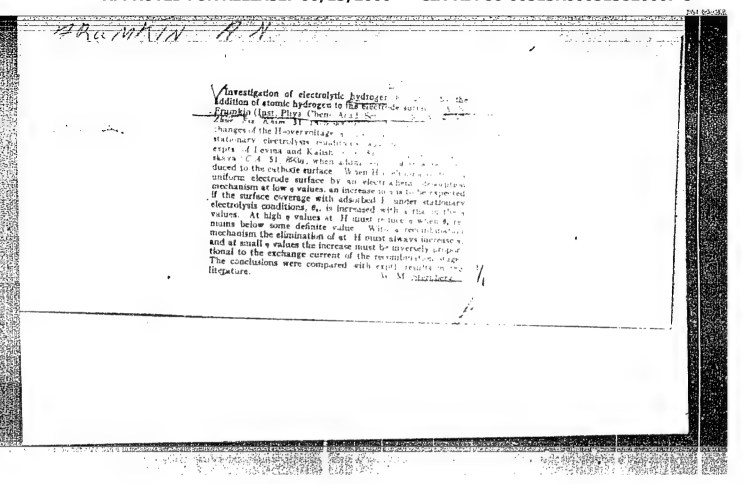
USSR/Physical Chemistry - Electrochemistry.

B-12

Abs Jour : Ref Zhur - Khimiya, No 1, 1958, 560

surface tension lowering cation. A detailed thermodynamic analysis was carried out; it showed that this conclusion did not disagree with the conclusion concerning the adsorption of atomic Tl on the boundary between the Tl amalgams and solutions not containing Tl ions (Frumkin A.N., Gorodetskaya A.V., Z. phys. Chem., 1928, 136, 451). It was assumed that not only simple, but also complex Tl ions can participitate in the adsorption process.

Card 2/2



IOFA, Z.A.; FRUNKIN, A.N.; MAZNICHENKO, E.A.

Effect of the nature of cations on the rate of hydrogen separation from alkaline solutions [with summary in English]. Zhur.fiz.khim. 31 no.9:2042-2051 S '57. (MIRA 11:1)

1. Moskovskiy gosudarstvenny universitet im. M.V. Lomonosova. (Cations) (Hydrogen) (Solution (Chemistry))

FRUNKIN, A.N. 20-4-33/60 Frumkin, A.N., Academician Damaskin, B.B., Nikolayeva-Fedorovich, N.V. **AU THORS** The Super-equivalent Adsorption of Cations on a Negatively Charged Mercury Surface. TITLE (Sverkhekvivalentnaya adsorbtsiya kationov na otritsatel'no zaryazhennoy poverkhnosti rtuti.) Doklady Akademii Nauk SSSR, 1957, Vol. 115, Nr 4, PERIODICAL pp. 751-754 (USSR) In a demonstration of electrocapillarity it is usually assumed that among the anorganic ions only the anions ABSTRACT possess a specific adsorbability; that the concentration of the anorganic cations in an electric double layer is only determined by the quantity of their charges. But there exist published data which point to the inaccuracy of such a conclusion. A direct conclusion on the different adsorbability of the cations of alkaline metals in solutions of 0,1 N - chlorides becomes clear from Grahame's paper who determined the precise values of the differential capacity of the latter. His conclusions are in bad agreement with experimental data. In order to avoid contradictions in the interpretation of test results, it is simpler to assume a certain, although not large, cation adsorption CARD 1/5

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20-4-33/60

The Super-Equivalent Adsorption of Cations on a Negatively Charged Mercury Surface.

with a larger radius. The study of the temperature coefficient of the reaction of electric anion reduction leads to the same conclusion. Measurements of the differential capacity in 0,01 N- chloride solutions of alkaline metals were performed, in order to check the guess of the specific cation adsorption. A pendent mercury drop served as test electrode. The obtained results are given in tab. 1 A. The curves concern the differential capacity in 0,01 Nsolutions of lithium, potassium and cesium chlorides. A difference in the capacity for cations with varius radii is here not only observed in the case of considerable negative potentials, but also close to the zero-charge point where a minimum occurs on the curves. A difference in the capacity values of lithium and cesium chloride solutions furnishes an additional indication concerning the specific adsorption of the cesium cation.

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The Super-Equivalent Adsorption of Cations on a Negatively Charged Mercury Surface.

This conclusion may, however, not be considered as unequivocal, sinds the data are in such solutions complicated by the C1- adsorption. The most convincing data concerning the super-equivalent cation adsorption may be obtained by measurements of the differential capacity on negatively-charged electrode surfaces in the presence of such an anion, as e.g. J-, whose adsorption in the surface layer markedly influences the value of the differential capacity. Such measurements are performed by the authors in 0,1 N solutions of

NaCl, NaJ, CaCl, CaJ, as well as in 1,1 N solutions

of

KC1, 1 N KJ + 0,1 N KC1, 1 N KC₁ + 0,1 N LaCl₅

and

1 N KJ + 0,1 N LaCl 3

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The data are given in fig. 1 B and 2 A. From them follows that in the case of sufficiently negative polarizations

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The Super-Equivalent Adsorption of Cations on a Negatively Charged Mercury Surface.

the iodine anion stops to exert an influence, and the values of the differential capacity for the corresponding chlorides and iodides are in agreement. In the case of still stronger negative potentials the espacity values are only determined by the extions present in the solution. But the potentials at whom the difference between the eapacity values of the corresponding chlorides and iodides disappears are not eugal and depend on the nature of the eation. As may be seen from fig. 1 B and 2 A, in the case of morehegative potentials the capacity in the presence of Cs+ or La+++ in connection with the penetration of amions into the electric double layer suddenly increases as compared to Ma+ or K+. The fact that the superequivalent adsorption of eations may lead to the penetration into the surface layer also in the ease of negative potentials, becomes especially obvious in the adsorption of organic eations. Fig. 2 B gives data concerning the dependence of the differential espacity in 1 N solutions of ECl, KBr and KJ in the presence of 10-3 N(CAHQ)A 280A.

CARD 4/5

CARD 5/5

APPROVED FOR RELEASE: 06/13/2000

CIA-RDP86-00513R000513820007

Frunkin, A.N. 20-1-27/12 Frumkin, A. N., Academician AUTHOR: On the Difference of the Potentials and the Accumulation of the Components of the Solution in the Diffusion Layer TITLE: During Steady Electrolysis (O raznosti potentsialov i nakoplenii sostavnykh chastey rastvora v diffuzionnom sloye

pri statsionarnom elektrolize)

Doklady AN SSSR, 1957, Vol. 117, Nr 1, pp. 102-105 (USSR) PERIODICAL:

The present report attracts attention to some peculiarities of the distribution of concentration and the potential in a ABSTRACT: diffusion layer during steady electrolysis. The author first investigates the case of the steady electrolysis of a binary

electrolyte with any concentration which consists of a

discharging cation with the valence n and a not discharging anion with the valence n. For the difference of the potentials between a point on the surface of the

electrode and a point in the center of the solution it

holds that $= (RT/n_2F)\ln(a_2''/a_2').$

However, the value of a"/a; cannot be determined from this formula, because not Φ , but only Φ + Δ_{∇} can be determined

Card 1/3

On the Difference of the Potentials and the Accumulation 20-1-27/42 of the Components of the Solution in the Diffusion Layer During Steady Electrolysis

 Δ_{ϕ} denotes the modification of the by measuring. Here potential difference ϕ between electrode and solution as a result of the change of concentration in the layer near the electrode. Next, the expression for Φ + $\Delta \phi$ which corresponds to the above formula, is written down. The measurable quantity Φ + Δ_{Ψ} is expressed not by the average but by the individual activities of the individual ions. It would be of great interest to apply the equation for $\P + \Delta \varphi$ the concentrated solutions, as e.g. acids. The author first investigates the case of a diluted acid which contains a cation 1, discharging on the cathode with the valence n and the concentration c₁ (in equivalents per cm3) as well as a not discharging anion 2 and a not discharging cation 3 with the valences n2 and n3 and the concentrations c2 and c3. Because of the condition of electric neutrality it holds that c₁ + c₂ = c₂. Next, relations between the concentrations of the not discharging ions on the surface of the electrode (c_2'', c_3'') and in the interior of the

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On the Difference of the Potentials and the Accumulation 20-1-27/42. of the Components of the Solution in the Diffusion Layer During Steady Electrolysis

solution outside the double layer during steady electrolysis are given. At $c_1^{\prime}/c_3^{\prime}=10^4$, $n_3=3$ and $n_2=1$ the degree of enrichment of the not dissolving cation is 10^3 . The practical applicability of such an enrichment process is, however, limited. However, the analytical applications of such an enrichment are real. In conclusion the author illustrates the relations between the total and the "ohmic" decrease of the potential in a diffusion layer on the basis of an electrolyte with three ions. There are 3 references, 2 of which are Slavic.

SUBMITTED: July 30, 1957

AVAILABLE: Library of Congress

Card 3/3

FRUMKIN, B.K.

AUTHOR:

Frumkin, A.N., Academician

25-1-8/48

TITLE:

Fuel Element (Toplivnyy element)

PERIODICAL:

Nauka i Zhizn', 1958, # 1, p 17 (USSR)

ABSTRACT:

The author, one of the leading Soviet electrochemists, recommends that research on the fuel element be intensified, e.g. the composition of highly efficient high temperature gas elements be studied and experiments with high temperature elements which make use of solid electrolytes with good conducting qualities be carried out. The discovery of such an electrolyte, where the transmission of current is carried out completely by oxygen anions, would represent

a remarkable achievement in this field of science.

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Library of Congress

Card 1/1

FRUMKIN, A.N., akademik; LUKOVTSEV, P.D., doktor khim. nauk

Present state of knowledge of the mechanism of electrode processes.

Khim. nauka i prom. 3 no.4:410-417 '58. (MIRA 11:10)

(Chemical reaction, Rate of) (Electrochemistry)

APPROVED FOR RELEASE: 06/13/2000 CIA-RDP86-00513R000513820007-5"

FRUMKIN, A.M.

AUTHOR:

Hone given

62-58-4-30/32

TITLE:

Anniversary Session of the Department

for Chemical Sciences of the AS USSR on October 30 and 31, 1957, and General Meeting of the Department for Chemical Sciences on lecember 19 and 20, 1957 (Yubileynaya sessiya otdeleniya khimicheskikh nauk Akademii nauk SSSR ot 30-31 oktyabrya 1957 g.i obshcheye sobraniye otdeleniya khimiches-

kikh nauk 19-20 dekabrya 1957 5)

PERIODICAL:

Izvestiya Akademii Hauk SSSR, Otdeleniye Khimicheskikh Nauk,

1958, Hr 4, pp. 521 - 524 (USSR)

ABSTRACT:

On the occasion of the 4oth anniversary of the October Revolution a reunion meeting of the Department for Chemical Sciences of the AS USSR took place. In his opening speech N. N. Semenov pointed out the outstanding succes of the USSR in the field of sciences especially in that of chemistry. Scientific lectures of the sessions were held by the following scientists, as was mentioned already earlier: Knunyants, Member, Academy of Sciences, and A. V. Fokin on the "Nitra-

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tion of Fluorofines", A. L. Midzhoyan, Member, AS Armenian SSR,

62-58-4-30/32

Anniversary Session of the Department for Chemical Sciences of the AS USOR on October 30 and 31,1957, and General Meeting of the Department for Chemical Sciences on December 19 and 20, 1957.

on the "Investigations in the Field of the Synthesis of Physiologically Active Compounds", R. Kh. Freydlina, Doctor of Chemical Sciences, reported on the "Investigation of the Telomerization Reaction and the Reaction of the Synthesis on the Basis of Telomers" (Reference 2). B. A. Dolgoplosk, Doctor of Chemical Sciences, spoke on the "Generation of Free Radicals in Solutions and Their Reactions in Model Systems," A. M. Frumkin, Member of the Academy of Sciences, reported on "Some General Problems of Electrochemical Kinetics and the Theory of Ion Reactions" (Reference 4), A. V. Kiselev, Doctor of Chemical Sciences (Reference 5) spoke on "Some Problems of Adsorption Theory", N. M. Emanuel (Reference 6), Doctor of Chemical Sciences, reported on "New Problems in the Field of Chain Reactions", V. L. Tal'rone, Candidate of Chemical Sciences, spoke on mass-spectroscopic investigations of ion-and radical reactions, A. P. Rebinder, Member,

Card 2/4

62-56-4-30/32

Anniversary Session of the Department for Chemical Sciences of the AS USSR on October 30 and 31, 1957, and General Meeting of the Department for Chemical Sciences On December 19 and 20, 1957

Academy of Sciences, drew conclusions with regard to the development of physico-chemical mechanics (Reference 7).

I. V. Tananayev, Corresponding Member of the AS USSR, gave new data on the chemistry of some rare elements, D. I. Ryabchikov and others spoke on the "Problems of the Chemistry of Rare Earth Elements"; the final lecture was that of V. A. Sokolov, Doctor of Chemical Sciences, on the "Calorimetric Measurements at High Temperatures".

General Regular Meeting of the Department for Chemistry of the AS USSR (December 19 - 20,1957):A. I. Brodskiy, Corresponding Member, AS USSR, spoke on the "Investigation of Some Reactions of Peroxides and Peracids of Hydrogen by Means of the Isotopic Method", M. M. Shemyakin, Corresponding Member, AS USSR, spoke on the "Use of N15 for the Explanation of the Mechanism of Some Organic Reactions", O. A. Reutov, Doctor of Chemical Sciences, reported on the "Investigation of the

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62-58-4-30/32

Anniversary Session of the Department for Chemical Sciences of the AS USUR on October 30 and 31,1957, and General Meeting of the Department for Chemical Sciences On December 19 and 20,1957

Electrophil and Homolytical Reactions of the Substitution in the Carbon Atom by Means of the Method of Isotope Exchange", I. P. Alimarin, Corresponding Member, AS USSE, reported on new methods of determination of the division of rare elements using organic derivative sulfuric-, selenicand telluric acids, V. G. Levich, Doctor of Chemical Sciences, reported on the "Diffusion Kinetics of Heterogenous Chemical Reactions in mobile Liquids". There are 8 references, all of which are Soviet.

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1. Chemical industry-USSR

Card 4/4

"APPROVED FOR RELEASE: 06/13/2000

CIA-RDP86-00513R000513820007-5

FRUHKIN, A.N.

76-1-24/32

AUTHORS:

Frumkin, A. N., Polyanovskaya, N. S.

TITLE:

Electrocapillary Phenomena in Salt Solutions of Thallium and Cadmium II. (Elektrokapillyarnyye yavleniya v rastvorakh soley talliya

i kadmiya. II.)

PERIODICAL:

Zhurnal Fizicheskoy Khimii, 1958, Vol. 32, Nr 1, pp.157-163 (USSR)

ABSTRACT:

Anomalous electrocapillary curves in thallium salt solutions. In the earlier works of the authors (reference 1) experimental results of the electrocapillary mercuric curves of solutions with thallium salts were given. The measurements were continued up to the potential φ =-0,45 with a normal calonel electrode. By this, the current passage through the solution in the capillary of the capillary-electrometer could be determined. On occasion of a control measurement carried out according to the method described in reference surement carried out according to the method described in reference with lightly acidulated 0,2 N TINO, + 0,8N KNO, it was shown that these measurements can be continued up to more negative potentials (N - normal). By this, a capillary curve of a strange form tials (N - normal). By this, a capillary curve of a strange form tials (N - normal) curves belonging to different systems thally two electrocapillary curves belonging to different systems are measured in experiment. The curve situated at the left is, as this is shown in the first part of this work, the electrocapillary

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76-1-24/32

Electrocapillary Phenomena in Salt Solutions of Thallium and Cadmium. II.

mercury curve of the thallium salt solution. Furthermore, it is shown that the right part of the two-humped curve shows the electrocapillary curve of the Tl-analgam with constant composition in a \sim 0.9N KNO, solution. It is shown that at $\gamma < -0.582$ with an accuracy of up to 10 % the composition of the amalgam forming in the capillary of the electrometer remains constant, whilst the concentration of the Tl ions decreases at least by the tenfold, i.e. up to 0,02 N. At ψ = -0,642 the boundary value of c_{Tl} (concentration of thallium in amalgam at the boundary with the solution) is obtained with an accuracy of up to 1 %, whilst the content to 0,002N. (concentration at the boundary with mercury) decreases up to 0,002N. The descending branch of the electrocapillary curve of the Tl-amalgam is near to that of the N KNO3, except it is displace somewhat to the negative side. In the reference 2 of one of the authors it is shown that the boundary-stress must increase by ~ 0.5 dyn/cm in a salution of equal composition at potentials being more nega- φ = -1,1 on occasion of the conversion from mercury to 0,45 % of amalgam. This fact corresponds to an average displacement of the descending branch by 3,5 mV in the direction towards the more negative potentials. This displacement is subjected to two further effects. The conditions for the arising of these effects will become clear, if the potential decrease and the quantity of

Card 2/4

76-1-24/32 1 Gadainm. II.

Electrocapillary Phenomena in Salt Solutions of Phallium and Gadainm. II.

the concentration variations produced on occasion of the current passage through the capillary are determined. If all these effects are summed up it can be seen that the descending branch of the electrocapillary curve 0,2H T1NO3 + 0,8N KNO3 in comparison to the same branch of N KNO $_{\rm x}$ solution must displace itself by $\sim 2,7+2,7+$ + 3,5 = 9 mV to the Regative side. The displacement observed on occasion of the experiment amounted to about 10 aV. The electrocapillary phenomena in cadmium salt solution. As it was shown by A. H. Frumkin and F. Servis (reference 3) the point of the zero-charge of the cadmium amalgam in KCl-, KBr- and KJ-solutions displace itself, as in the case of thallium salts into the direction of the more negative potentials. This was also confirmed by the neasurements of the electrocapillary curves of the analgam by A. V. Gorodetskaya. It is shown that the total curve obtained on occasion of the measurement of the dependence of the boundarystress of mercury upon the polarization in the 2,0% ABr + 0,5% CdBr2-solution distinctly consists of two electrocapillary curves. The left one is to be considered as the general electrocapillary curve of mercury in the solution mentioned, wailst the right curve has to be taken for a electrocapillary curve of cadmium aumlgam in the KBr-solution. Analogous results were obtained with the 0,8N KBr+

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CIA-RDP86-00513R000513820007-5 "APPROVED FOR RELEASE: 06/13/2000

76-1-24/32

Electrocapillary Phenomena in Salt Solutions of Thallium and Cadmiun. II.

+ 0,2N CdBr₂-solution, whilst in the case of the 0,0N Ma₂SO₄ + 0,2N CdSO₄-solution a salient point in the total $\mathcal{E} = \varphi$ -curve couldnot be found. It is assumed that in this system the surface activity, as well of the Cd²-ion in the solution, as of the Cd discoluted in the course of the Cd²-ion in the solution. dissolved in mercury, is much too small. There are j figures, and 6 references, all of which are Slavic.

ASSOCIATION:

AS USSR. Institute of Physical Chemistry. Hoscow (Akademiya nauk SSSR. Institut fizicheskoy khimii. Moskva)

SUBMITTED:

November 5, 1956

AVAILABLE:

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CIA-RDP86-00513R000513820007-5 "APPROVED FOR RELEASE: 06/13/2000

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Frunkin, A. N., Academician

20-3-32/59

AUTHORS:

TITLE:

Tedoradze, G. A.

The Ionisation Kinetics of Eolecular Chlorine

(Kinetika ionizatsii molekulyarnogo khlora)

PERIODICAL:

Doklady AN SSSR, 1958, Vol. 118, Nr 3, pp. 530-533 (USSR)

ABSTRACT:

The authors investigated the kinetics of the electric reduction of chlorine on a rotating disk-shaped platinum 2 electrode. The surface area of the electrode was 0,78 mm². The chlorine was produced by electrolysis of 15- per cent HCl and washed by letting it pass through the solution in test. The electrode to be investigated was activated before the experiment. The performance of the experiment is described shortly. At not very high excess voltages the amperage of the ionisation current for chlorine depends on the fact, wether the voltage changes from the more anodic potentials to the more cathodic ones or vice-versa. A limiting value of the amperage of the ionisation current must exist, which is determined by the dissociation velocity of the chlorine molecules. Besides, no linear dependence between the excess voltage and the logarithm of the cathode

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current is obtained. The amperage of the cathode current in case of given excess voltage cannot depend on the con- : 1. centration of the chlorine ions [Cl-]. For the potential of the electrode and for the excess voltage formulae are given. In case of validity of a certain assumption given here in detail a linear dependence must be valid between the logarithm of the corrent density and the potential (or the overvoltage). This conclusion is also proved by the experiment. The independence of the current density on [C1-] in case of given potential of the electrode proves the irreversibility of the state of self-sustained ionisation. The results of the experiments fit well into the score of the theory, if the following is assumed: The process of the ionisation of Cl2 passes the following two states: Cl + e = Cl adsorbed + Cl; Cl adsorbed + e = cl apart from the state of the diffusion of Cl2. The first one of these both reactions is irreversible in case of sufficiently high excess voltage. There are 4 figures and 4 references, all of which are Slavic.

ASSOCIATION:

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20-3-32/59

SUBMITTED:

September 25, 1957

AVAILABLE:

Library of Congress

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20-119-2-36/60

AUTHOR:

Frumkin, A. N., Member, Academy of Sciences, USSR

TITLE:

On the Stoichiometric Number of the Reaction of Electrochemical Hydrogen Desorption (O stekhiometricheskom chisle

reaktsii elektrokhimicheskoy desorbtsii vodoroda)

PERIODICAL:

Doklady Akademii Nauk SSSR, 1958, Vol. 119, Nr 2

pp 318 - 321 (USSR)

ABSTRACT:

First several previous papers dealing with the same subject are mentioned, discussing the applicability of a criterion proposed by J. Horiuti and Yukasima (Reference 8) concerning the explanation of the mechanism and of the nature of the slow stage of the transition of hydrogen ions into molecular

hydrogen

 $(2H^+ + 2e^- + H_2)$.

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At present usually the stoichiometric number ν which is equal to $2/\lambda$ is used as criterion. Here ν denotes a number

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characterizing the process. This number ν expresses how many times the slowest stage must be repeated in oder to pass the whole process once. When the surface of the electrode is homogenous and when the displacement of $\mu_{\rm adsorbed}$ on the surface of the electrode is not connected with a noticeable activation energy the amounts of direct and inverse reaction velocities ν_1 and ν_2 of the stage discharge-ionization

and of the direct and inverse reaction velocities $\ v_{3}$ and v_{4} of the stage of electrochemical desorption-adsorption

 $H_{adsorbed} + H^{\dagger} + e^{-} \longleftrightarrow H_{2}$ can be represented by the here given equations, The further

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course of the calculation is followed step by step and the results are mentioned in detail:

 $\nu = 1 + 1/2\gamma = 1 + i_{OT}/2i_{OTT}$

Here $\cdot i_{OI}$ denotes the exchange current in the stage discharge-ionization and i_{OII} denotes the exchange current in the stage desorption-adsorption. Only with $i_{OI} \ll i_{OII} = \nu = 1$ holds valid but with $i_{OI} \gg i_{OII} = \nu$ can become arbitrary great. According to the equations found here the existence of a considerable exchange current i_{OII} with small i_{OI} secures a great value of the total exchange current, but not a small polarizability of the electrode. The calculations carried out for ν do not furnish a correct idea on the values of ν given in references. The reasons for this are shortly

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discussed.With $\gamma \ll 1$ the extrapolated amperage is not equal to the real amperage. On certain here given conditions the value of $\mathcal V$! determined from the extrapolated power of the exchange current is equal to unity. The same value for $\mathcal V$! is obtained when the anode part of the curve and not the cathode part is used for extrapolation. There are 13 references,7 of which are Soviet.

SUBMITTED:

December 24, 1957

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AUTHORS:

Damaskin, B. B., Nikolayeva-Fedorovich, SOV/20-121-1-36/55

N.V., Frumkin, A. N., Member, Academy of Sciences, USSR

TITLE:

On the Adsorption of the Ion Cs⁺ on the Surface of a Mercury Electrode (Ob adsorptsii iona Cs⁺ na poverkhnosti rtutnogo

elektroda)

PERIODICAL:

Doklady Akademii nauk SSSR, 1958, Vol. 121, Nr 1,

pp. 129 - 132 (USSR)

ABSTRACT:

The authors performed their measurements at solutions of sodium fluoride and cesium fluoride to avoid complications in the investigation of the superequivalent (sverkhekvivalent-noy) cation adsorption. The authors determined the differential capacitance by means of an impendance bridge at a dropping-mercury electrode in solutions of NaF and CsF of concentrations of 0,9 N, 0,1 N, and 0,01 N as well as in solutions of NaJ and CsJ. The data found are given, in 2 diagrams. The capacitance of the double layer in solutions of cesium fluoride is higher than in solutions of sodium fluoride. Numerical data are given as well. According to the obtained data a cesium cation is

more adsorbed on the mercury surface (on which a fluorine

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is adsorbed) than a sodium cation. A conclusion on the superequivalent adsorption of Cs⁺ can also be made by comparing the corresponding curves of the capacitance of the fluorides and iodides of sodium and cesium. These data on the superequivalent adsorption of cesium were also proved by electrocapillary measurements in 0,1 N solutions of NaF, NaJ, CsF, and CsJ. An important fact is also the displacement of the maximum on the electrocapillary curve from -0,471 in the case of NaF to the value of -0,468 for CsF and from -0,815 for NaJ to -0,833 for CsJ. There are 3 figures, 1 table, and 9 references, 3 of which are Soviet.

SUBMITTED:

May 8, 1958

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On the Adsorption of the Ion Cs⁺ on the Surface of a SOV/20-121-1-36/55

1. Cesium ions--Adsorption 2. Mercury electrodes--Adsorptive properties

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